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Genome-wide association defines more than thirty distinct susceptibility loci for Crohn's disease

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Abstract

Several new risk factors for Crohn's disease have been identified in recent genome-wide association studies. To advance gene discovery further we have combined the data from three studies (a total of 3,230 cases and 4,829 controls) and performed replication in 3,664 independent cases with a mixture of population-based and family-based controls. The results strongly confirm 11 previously reported loci and provide genome-wide significant evidence for 21 new loci, including the regions containing *STAT3*, *JAK2*, *ICOSLG*, *CDKAL1*, and *ITLNI*. The expanded molecular understanding of the basis of disease offers promise for informed therapeutic development.

The first genome-wide association studies (GWAS) have identified many common variants associated with complex diseases, and have rapidly expanded our knowledge of the genetic architecture of these traits. Progress in Crohn's disease (CD), a common idiopathic inflammatory bowel disease (IBD) with high heritability ($\lambda_s \sim 20\text{--}35$), has been especially striking, with recent GWAS publications increasing the number of confirmed associated loci from two to more than ten¹. The results have identified new pathogenic mechanisms of IBD and promise to advance fundamentally our understanding of CD biology. These recent discoveries highlight, for instance, the key importance of autophagy and innate immunity²⁻⁵ as determinants of the dysregulated host-bacterial interactions implicated in disease pathogenesis. Furthermore, genetic associations have been shown to be shared between CD and other auto-inflammatory conditions – for example, *IL23R* variants⁶ are also associated with psoriasis⁷ and ankylosing spondylitis⁸, and *PTPN2* variants with type 1 diabetes^{3,5}. As in other complex diseases, restricted sample sizes have resulted in early CD studies focusing on only the strongest effects, which turn out to explain only a fraction of the heritability of disease.

We recently published three separate GWA scans for CD in European-derived populations – the details of which are shown in Table 1^{4,5,9}. Motivated by the need for larger datasets to improve power to detect loci of modest effect, we carried out a genome-wide meta-analysis from our three CD scans. These analyses, together with a replication study in an equivalently sized, independent panel, have enabled us to identify at genome-wide levels of significance 21 novel Crohn's disease susceptibility genes and loci. This brings the total number of independent loci conclusively associated with Crohn's disease to more than 30 and provides unprecedented insight into both CD pathogenesis as well as the general genetic architecture of a multifactorial disease.

Results

Meta-analysis of three genome-wide association scans

The combined GWAS study samples (Table 1) consisted of 3,230 cases and 4,829 controls, all of European descent. While the individual scans did identify new risk factors, they were only well-powered to discover common alleles with odds-ratios (ORs) above 1.3 (in the case of the WTCCC) or 1.5 (the smaller two scans, Figure 1). By contrast, the combined sample has 74% power at an OR of 1.2, allowing evaluation of the role of alleles with smaller effect sizes for the first time. As two different genotyping technologies were used in the constituent scans, we utilized recently developed imputation^{10,11} methods to assess association across all three studies at 635,547 SNPs contained on one or both platforms. A quantile-quantile (Q-Q) plot of the primary meta-statistic (single SNP Z-scores, Figure 2) shows a striking excess of significant associations, well beyond what would be attributable to the modest overall distributional inflation (genomic control $\lambda < 1.16$). Despite the large sample size, the overall inflation is modest because (1) each group had separately tested for evidence of population stratification, and the meta-analysis used a test that combined the results from each study (rather than mixing the raw data and compromising the case-control matching of each study), and (2) imputation was done on all samples ignoring case status and thus would not introduce artifactual differences between cases and controls¹².

We focus our attention in this study specifically on the 526 SNPs from 74 distinct genomic loci which were associated with $p < 5 \times 10^{-5}$ – more than 7 times the number of SNPs expected by chance even after correction for the modest overall inflation detected. This threshold for follow-up is not meant to imply that there are no genuine associations among SNPs with less significant association in the meta-analysis, but rather reflects a practical desire to prioritize as many true positives as possible for immediate replication. Eleven associations previously replicated and established at genome-wide significance levels (Methods, Table 2), including

both “historical” associations at *NOD2*^{13,14} and 5q31 (IBD5)¹⁵ as well as recent replicated findings from individual GWA scans such as *IL23R*, *ATG16L1*, *IRGM*, *TNFSF15* and *PTPN22*^{6,16}, were among the 74 regions represented in this tail of the distribution of association statistics. Even after removing all SNPs in LD with these eleven loci, however, there continued to be a substantial excess of associated alleles beyond that which would be expected by chance (Figure 2).

Replication of 21 new loci

As these 74 regions included the 11 already reported as independently replicated and meeting genome-wide significance thresholds, this replication experiment effectively explored 63 putative associations in novel regions with 11 positive controls (Supplementary Table 1). To identify the true risk factors from these 63 regions, we undertook a replication study involving a total of 2,325 additional Crohn's disease cases and 1,809 controls alongside an independent family-based dataset of 1,339 parent-parent-affected offspring trios.

Results (significance levels and odds ratios) for strongly replicating loci, including all positive controls, are presented in Table 2. The distribution of Z-scores from the 63 putative regions shows a dramatic departure from the null distribution (Figure 3) with 19 novel regions showing significant replication ($p < 0.0008$ – a value of $0.05/63$ representing a conservative threshold expected to be exceeded only once by chance in 20 such replication experiments). SNPs on chromosome 19p13 (replication $p = 0.00347$, combined $p = 2.12 \times 10^{-9}$) and in the MHC (replication $p = 0.006$, combined $p = 5.2 \times 10^{-9}$ - suspected but not previously conclusively established in Crohn's disease) did not reach this conservative threshold, but so convincingly satisfy proposed thresholds for genome-wide significance ($p < 5 \times 10^{-8}$, Methods) that we propose these as the 20th and 21st additional Crohn's disease associated loci defined here. A further 8 of the 42 remaining loci showed nominal replication (Table 3).

It is possible that extreme population substructure in the replication sample could give rise to such a striking excess of hits. While unlikely, this was directly evaluated by the large family-based component of the replication study. Odds ratio estimates from the TDT analysis of the North American, French and Belgian families alone are consistent with those from the UK and Belgian case/control samples (Tables 2 & 3), with all 21 newly defined loci showing odds ratios in the same direction of association with the original scan in the family-based component (and nearly half showing greater OR than in the case-control arm). Importantly, none of the significantly or nominally replicating loci show significant evidence for heterogeneity (across studies or between family-based and population-based arms) when corrected for the number of tests performed. This independent family based evidence (Supplementary Table 6) confirms these alleles constitute true Crohn's disease loci.

For this newly expanded set of 32 unequivocally associated loci, we assessed whether there was evidence of significant pairwise interactions which could add further to the overall variance in liability explained by this set of loci. We performed a case-only analysis of the 3,664 cases in the replication study and observed no interactions that withstood a correction for the number of tests performed (Supplementary Table 2).

Deciphering the genetic architecture of CD

The contributions of the 32 loci to disease risk were computed using a standard liability threshold model and are displayed as a histogram of individual variances (Figure 4). The observations from this variance analysis that many loci were detected for which the current study had low power, and that only a minority of the variance in risk is explained by these 32 loci, suggest that many additional loci are yet to be identified. This is reinforced by the additional 8 nominal replications (Table 3) where only 2 or 3 would be expected by chance,

and by the continued excess of small p values when these 40 total regions are removed (Figure 2).

While recognizing that fine-mapping is required to identify specific causal variants, we performed a series of analyses to gain some general insight into the CD associations. We first queried HapMap to discover any instances where a non-synonymous SNP (nsSNP) was correlated ($r^2 > 0.5$) to the most associated variant discovered in this study. Accepting that HapMap is not a complete catalogue of nsSNPs, but including four loci where fine-mapping has identified coding variants, just 9 of the 32 genomewide significant associations were correlated with a known nsSNP (Supplementary Table 3). To explore whether any of the associations reflect a cis-acting regulatory effect on a nearby gene, we evaluated genotype-expression correlation using the panel of 400 lymphoblastoid cell lines described by Dixon et al.¹⁷. From all genes within 250 kb of the LD-based intervals defined in Table 2 and 3, five correlations between expression of a nearby gene and a CD-associated variant were identified (LOD > 2) (Supplementary Table 4). This was far in excess of chance ($p \sim 0.001$) (Supplementary Figure 1) and suggests that regulatory variation also contributes to the genetic architecture identified.

Discussion

Genome-wide association studies provide a systematic assessment of the contribution of common variation to disease pathogenesis. A limiting factor is often the size of the case-control dataset, and hence the power to detect any but the most strongly associated loci. Meta-analysis of existing data provides an obvious potential solution. As Figure 1 demonstrates, our expectation was that the additional power of the combined dataset would result in the identification of a substantially larger number of readily replicating associations than were derived from any of the smaller, constituent datasets. However, the paradigm of exploring common genetic variation with similar effects across studies (in this case all of European descent) needs testing before its results can be accepted as valid.

On the validity of the method our results are substantially reassuring. All 11 previously confirmed CD susceptibility loci were strongly replicated both in the meta-analysis and follow-up experiment. These include the two widely replicated findings from studies published in 2001¹³⁻¹⁵ as well as all of the compelling findings from individual GWAS (Table 2 a). Significantly, we have also identified and replicated 21 new CD susceptibility loci. Using a conservative threshold for significance (only 1 such region would be expected by chance in 20 such experiments), the loci with clear evidence for association in the replication panel include a very high proportion of those showing strongest signals in the meta-analysis (Supplementary Table 1) – 9 of 9 previously unreported regions with $p < 5 \times 10^{-7}$ in the combined scan were replicated convincingly - emphasizing the validity of the meta-analysis results. Further emphasizing the robustness of these results, all 21 of these loci exceed a conservative genome-wide level of significance ($p < 5 \times 10^{-8}$) by a significant margin (all but two have $p < 5 \times 10^{-9}$) - and equivalent strength of association was observed in the family-based subset of our replication sample.

In keeping with other regions recently identified as associated with CD, the 21 new loci do not conform to any obvious pattern in terms of gene content. Thus, as shown in Table 2, some loci (defined by HapMap recombination hotspots flanking the set of correlated, associated variants) contain just a single gene, some contain many genes and others none. Clearly the first category provides the most immediate clues regarding pathogenic mechanisms. These genes are discussed briefly in Box 1, together with a number of genes which constitute striking candidates from regions with only a handful of transcripts. Included among these are compelling functional candidates such as *STAT3*, *JAK2* and *IL12B* while others, such as *CDKAL1* and *PTPN22*,

highlight potentially intriguing contrasts between genetic susceptibility to Crohn's disease and some other complex disorders (Box 1). It is noteworthy – and consistent with previous findings from CD and other complex diseases – that we did not find any strong evidence of deviation from the model of multiplicative (random) effects when we tested for gene-gene interactions among the 32 confirmed associations. This is in spite of the fact that some of these genes seem to affect the same or overlapping pathways.

For loci containing multiple genes or no genes the picture is less well defined. The identified paucity of correlation between associated SNPs and coding variation suggests that these loci may, in particular, benefit from eQTL (expression quantitative trait locus) analysis. This seeks correlation between genotype and expression patterns – bearing in mind that such functional relationships need not respect the specific boundaries of LD around the association. One of our groups previously reported an eQTL effect incriminating *PTGER4* at the 5p13 locus⁹. A striking outcome from our present analysis was at the established IBD5 locus¹⁵, where CD-associated SNPs were associated with decreased *SLC22A5* mRNA expression levels. While a SNP had previously been proposed as regulating *SLC22A5* transcriptional activity¹⁸, these data suggest for the first time that the most disease-associated variants in the IBD5 region, including a coding variant in neighboring *SLC22A4*, are the same variants most associated with *SLC22A5* expression. Equally striking, the most significant Crohn's disease associated eQTL reported here affects *ORMDL3* (LOD = 20) on chromosome 17 and SNPs in precisely the same region were recently shown to be strongly associated with childhood asthma.¹⁹ This suggests that the same polymorphisms might underlie susceptibility to both CD and asthma, possibly by perturbing *ORMDL3* expression.

The new loci that we have identified are of modest effect size, which is unsurprising given all loci with larger impact on disease risk were – as might be expected – discovered in the original scans. The small sizes of these effects explains the lack of overlap between linkage results in CD and these newly discovered loci (Supplementary Figure 2), with the possible exceptions of combined effects of multiple high ranking associations on chromosomes 5q and 6p. Indeed, the linkage evidence that led to the discovery of the IBD5 locus was very likely boosted by the nearby effects at *IL12B* and *IRGM*. As expected, the only gene conclusively discovered via linkage (*NOD2*) is one of two loci which stand well out from the remainder of the distribution of effect sizes (Figure 4). The other outlier, *IL23R*, illustrates an interesting characteristic of linkage – because (unlike *NOD2*) the most penetrant risk allele has very high frequency (93%), it is nearly invisible to linkage analysis despite the high OR; highly protective rare alleles are simply not present in multiplex affected families and thus do not influence allele sharing substantially.

Using a liability-threshold model, we estimate that the 32 loci identified to date explain about 10% of the overall variance in disease risk, which may be as much as a fifth of the genetic risk, given previous estimates of CD heritability of approximately 50%.²⁰ This observation is consistent with the fact that these loci collectively contribute only a factor of two to sibling relative risk (λ_s), and even this figure is dominated by the substantial contribution of *NOD2* variants. However, it should be emphasized that the full impact of the new loci cannot be determined until causal variants have been identified by directed sequencing and fine-mapping experiments. Until then the proportion of the variance in Crohn's disease risk explained must be measured from the confirmed SNPs, where association is due to LD with causal variants. Since multiple causal variants might exist at each locus (ranging in frequency from rare to common) our estimates of variance explained provide only a lower bound for the true contribution of each locus.

In conjunction with results from a very similar gene discovery effort in type 2 diabetes²¹, common lessons are beginning to emerge with respect to the genetic architecture of complex

traits. In each example, substantial increase in sample size achieved through meta-analysis has led to dramatic success in gene discovery. In all cases, this progress has revealed an underlying architecture consistent with many individually modest effects which conventional genetic linkage analysis, and even the largest individual genome-wide association studies, are not well powered to detect. Common variants explaining more than 1% of the genetic variance are rare, whereas well-powered studies have found dozens of variants contributing 0.1% of overall variance in liability. Perhaps surprisingly, neither we nor others have yet to document a substantial role for epistasis among these loci and a number of associated loci are conclusively mapped to regions with no currently annotated protein coding genes. Despite the considerable concordant success, a distinct minority of the overall heritability has been explained by these documented associations.

Since our study is well-powered to identify loci that explain $> 0.2\%$ of the overall variance, but the sum of such loci explains a relatively small fraction of the total, it seems likely that many loci with even more modest effect sizes remain undiscovered. Of particular note is the continued excess of associations outside of the regions studied here, as well as the nominal replication of an additional 8 loci, notably greater than expected by chance. Overall, the distribution of Z scores in the replication experiment is clearly skewed towards replication – only 11 of the 63 Z-scores in this replication experiment generate $Z < 0$. If only the 21 strongly confirmed loci were genuinely associated, half of the 42 remaining should end up with $Z < 0$. Indeed, observing 8 of the 42 remaining tests with $Z > 1.5$ is itself a highly significant observation ($p < 0.0001$). Although modest in terms of effect size, identification of such loci is likely to still provide important insights into pathogenic mechanisms, as biological importance need not be proportional to the statistical evidence for genetic association. Closer inspection of regions showing nominal association in the replication experiment reveals that a number of transcripts in these loci are of considerable interest, including *CCL2/CCL7*²², *IL18RAP*²³ and *GCKR*²⁴.

It is important to note that the generation of GWAS arrays used in the scans here did not offer complete genome coverage of common variation (additional loci may reside in poorly covered intervals) and did not address either rare SNPs or copy number variation effectively. Thus in spite of the wealth of new susceptibility genes and loci identified by the current study, it seems implausible that there are not more to be found – albeit very large datasets are likely to be required to achieve robust statistical support for them. With respect to the present findings, there is much work to be done in resequencing and fine mapping to identify causal variants. While we do not yet have a complete understanding of the genetic architecture of Crohn's disease, dramatic progress has now been made towards this goal - and with it the prospect of directed functional exploration of the pathways identified, insight into how risk alleles interact with environmental modifiers, and the hope of new avenues for treatment.

BOX 1

Noteworthy genes within loci newly implicated in Crohn's pathogenesis

- Chemokine receptor 6 (*CCR6*): encoding a member of the G protein-coupled chemokine receptor family, this homing receptor is expressed by immature dendritic cells and memory T cells and is important for B-cell differentiation and tissue specific migration of dendritic and T cells during epithelial inflammatory and immunological responses²⁵. The ligand of this receptor is macrophage inflammatory protein 3 alpha (MIP-3 alpha); both genes are expressed in granulomas of pulmonary sarcoid²⁶. Recent studies have also demonstrated that *CCR6*, *IL23R* and *RORγT* are selectively expressed by *IL-17* producing cells and *IFNγ* producing TH17/TH1 cells in CD²⁷.

- Interleukin ***IL12B***: encodes the p40 subunit which is a constituent of both heterodimeric interleukins IL-12 and IL-23²⁸. Association with CD was previously reported⁵ but not confirmed, and it is also known to be associated with psoriasis⁷. The key role of the IL12/IL23 pathway in chronic intestinal inflammation is supported by the association between *IL23R* and CD³ and strong functional evidence from mouse models of colitis²⁹⁻³².
- Signal transducer and activator of transcription 3 (***STAT3***) and Janus kinase 2 (***JAK2***): the JAK-STAT pathway is a focal point in signal transmission downstream of cytokine and growth factor signals from cell surface receptors to the nucleus to modify transcription of various genes, notably in hematopoietic cells. The present findings are particularly significant, given the role of both genes in *IL23R* signaling³³, and the central role *STAT3* in Th17 differentiation³⁴. However, *JAK2* or *STAT3* are also downstream of several other cytokines implicated in CD pathogenesis in addition to interleukin 23, highlighting the pathophysiologic complexity of these new associations. Further complexity is highlighted by the distinctly different roles of *STAT3* in innate versus adaptive immunity in murine colitis models: activation of *STAT3* in innate immune cells enhances mucosal barrier function whereas *STAT3* activation in T-cells exacerbates colitis.
- Leucine-rich repeat kinase 2 (***LRRK2***). This gene encodes a multi-domain protein expressed mainly in the cytoplasm of neurons, myeloid cells and monocytes, and mutations in *LRRK2* have been strongly associated with Parkinson's disease³⁵. A recent study reported the induction of autophagy by mutant *LRRK2*,⁴¹ which is of interest given the strong associations between CD and the autophagy genes *ATG16L1* and *IRGM*.²⁻⁵ The same locus also contains the gene ***MUC19***, which encodes a large protein with multiple serine/threonine-rich repeats characteristic of the mucin gene family. The mucin proteins are core components of the mucus layer which protects the intestinal epithelia from injury, and mucin-deficiency potentiates intestinal inflammation in mouse models of colitis³⁶.
- ***CDKAL1***: the protein encoded by this gene is poorly characterized, but *CDKAL1* is noteworthy for being recently confirmed as a type 2 diabetes susceptibility gene^{24,37-39}. In this study, we find that SNPs from the same intron of *CDKAL1* that shows association with T2D are associated with CD, but the associated alleles for the two diseases are not correlated with each other.
- Inducible T-cell co-stimulator ligand (***ICOSLG***): this co-stimulatory molecule is expressed on intestinal (and other) epithelial cells and may play a role in their antigen presentation to and regulation of mucosal T lymphocytes⁴⁰. Upon maturation, plasmacytoid dendritic cells express *ICOSLG* and drive the generation of IL-10 producing T regulatory cells⁴¹.
- Protein tyrosine phosphatase, non-receptor types 2 and 22 (***PTPN2* and *PTPN22***). Both of these genes are associated with other autoimmune and inflammatory diseases and the effect described here for *PTPN2* is similar to that previously described for type 1 diabetes (T1D)⁴². However, the association of *PTPN22* with CD, although mapping to the same coding variant (R602W) that is a risk factor for T1D and rheumatoid arthritis,^{43,44} is in the opposite direction, with the T1D and RA risk allele, 602W, offering protection from CD.
- Intelectin 1 (***ITLN1***) is known to be expressed in human small bowel and colon, and encodes a 120-kDa homotrimeric lectin recognizing galactofuranosyl residues found in cell walls of various microorganisms but not in mammals⁴⁵. Human intelectin-1 is structurally identical to the lactoferrin receptor (LFR), expressed

within the enterocyte brush border, and appears critical in membrane stabilization, preventing loss of digestive enzymes, and protecting the glycolipid microdomains from pathogens⁴⁶. In addition, intelectin expression is reported in Paneth cells in both mouse and pig small intestine, further pointing to a role in innate immunity.

Methods

Crohn's disease patients, controls, and GWAS

The meta-analysis was based on data from the 3 genome-wide scans of the NIDDK⁴, WTCCC⁵ and Belgian/French⁹ studies. Details of the numbers of cases and controls genotyped in the respective scans and of the genotyping platforms used are shown in Table 1, as are case/control and family cohorts genotyped in the replication study of the meta-analysis. Details of the ascertainment and characterization of these cohorts, as well as quality control procedures applied to the GWA datasets, were provided in the original scan and replication publications^{3, 4, 5, 6, 9}. Recruitment of study subjects was approved by local and national institutional review boards, and informed consent was obtained from all participants.

Imputation

Briefly, these methods rely on observed haplotype patterns in a set of reference data (the HapMap) and the actual genotype data from each project to make predictions (along with a measure of statistical certainty) at un-genotyped SNPs. We used the program MACH¹⁰ with the NIDDK and Belgian/French data, and IMPUTE¹¹ with the WTCCC data. Comparisons between the two algorithms yielded very similar results (data not shown). We imputed the superset of polymorphic markers which passed QC in the original scans^{4,5,9}. This set was comprised of SNPs on either the Affymetrix 500K only ($n = 350,507$), Illumina HumanHap300 version 1 only ($n = 238,935$), or both panels ($n = 46,105$) such that all association tests performed were at least partially based on observed genotype data.

Test for association, effect size estimation and interactions

Using the genotype probabilities (rather than best-guess genotypes) and empirical variances for imputed markers in the case and control tallies, we summarized the standard 1 d.f. allele-based test of association as a Z-score within each scan and combined scores across studies to produce a single meta-statistic for each SNP across all three datasets. Odds ratios were estimated separately in TDT samples and each case/control replication collection, and then combined and tested for heterogeneity.⁴⁷ Interaction tests were performed using the case-only epistasis test implemented in PLINK⁴⁸.

Critical regions

Given that most associations contain many correlated SNPs showing signal, we demarcated independent loci by first defining the set of HapMap SNPs with $r^2 > 0.5$ to the most significantly associated SNP. We then bounded the "critical region" by the flanking HapMap recombination hotspots which contained this set. These windows very likely contain the causal polymorphisms explaining the associations.

Replication

We defined loci to have been previously confirmed if an earlier study had both detected and replicated the association in independent samples and the association achieved $p < 5 \times 10^{-8}$ (recently proposed as an appropriate genome-wide significance level for GWAS⁴⁹). For replication genotyping, we selected the most significantly associated SNP from each region along with a second, correlated SNP with $p < 0.0001$ or a second assay on the opposite strand

in order to have a technical backup should the first fail genotyping (Supplementary Table 1). Replication genotyping for the putatively associated loci was performed using primer extension chemistry and mass spectrometric analysis (iPLEX, Sequenom) using Sequenom Genetics Services (N. American panel) and Genome Research Limited, Wellcome Trust Sanger Institute (UK panel), and using a custom-made Golden Gate assay on a Beadstation500 (Illumina), following the manufacturer's recommendations (Belgian/French panel). The more completely genotyped SNP of the two from each region was chosen to represent that regional association in analysis (if both were completely typed, the SNP that was more strongly associated in the scan was used). Samples with >10% missing data ($n = 267$ for Belgian/French data, 111 for the UK data and 8 for the N. American data; these samples are not included in the tallies for Table 1), as well as SNPs with >10% missing data or Hardy-Weinberg p value < 0.001 were excluded from this analysis.

Regional Annotation: eQTL analysis

Effects of SNPs in Tables 2 & 3 on expression levels of neighbouring genes was studied using transcriptome data from the ~400 lymphoblastoid cell lines described by Dixon et al.¹⁷. SNPs that were not genotyped on this panel ($n=14$) were replaced with a proxy with $r^2 > 0.95$ when possible ($n=12$). LOD scores > 2 for genes (probe average) located within 250 Kb of the corresponding LD windows were retrieved from <http://www.sph.umich.edu/csg/liang/asthma/>. To evaluate the significance of the findings with the CD associated SNPs, we compared the observed (i) number of genes yielding LOD scores > 2 , and (ii) sum of these LOD scores, with the corresponding frequency distributions for 1,000 randomly selected sets of 31 SNPs, matched for allele frequency (± 0.02) and gene context. Window sizes determined for associated SNPs were used for the matched simulated SNPs.

URL

Meta-analysis test statistics and allele frequencies for all SNPs are available at: <http://www.broad.mit.edu/~jcbarrret/ibd-meta/>

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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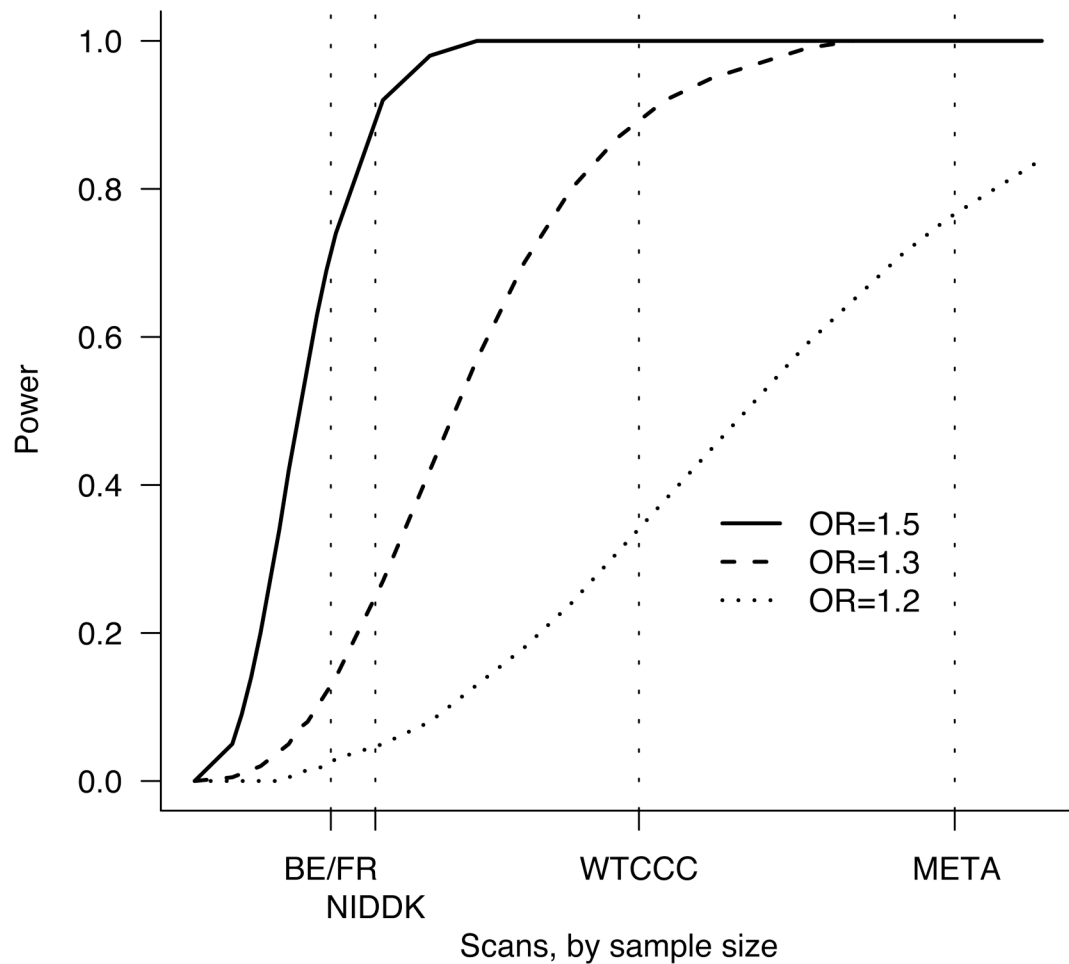


Figure 1.

Power to detect a genetic effect of various sizes (odds ratio 1.2, 1.3, 1.5) versus study sample size. Power is reported here as the probability (given a multiplicative model and risk allele frequency of 20%) of $p < 5 \times 10^{-5}$ in a scan – the value used to define regions for attempting replication in a larger sample set. Vertical dotted lines show the sample sizes for the three constituent scans and the meta-analysis. Relatively large effects are likely to be detected by any of these scans, whereas only the combined analysis is well powered to detect more modest effects.

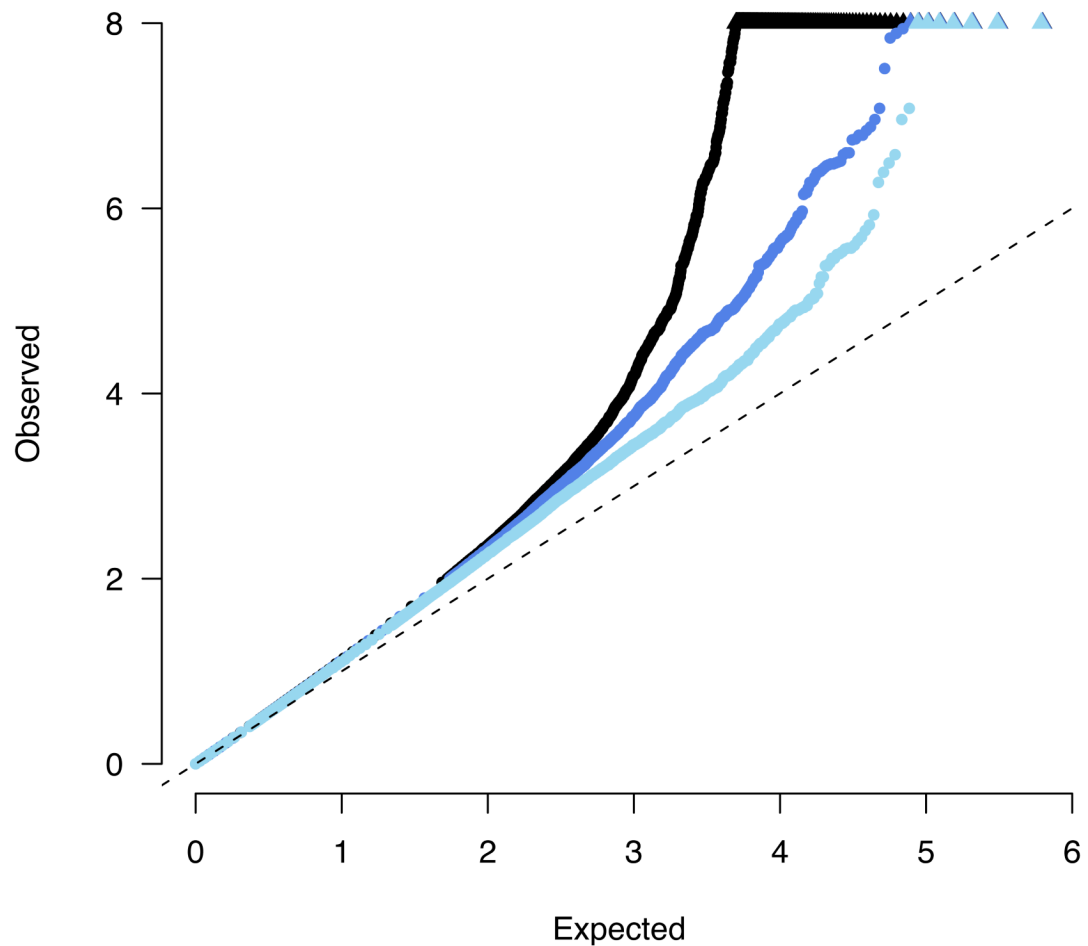


Figure 2.

A quantile-quantile plot of observed $-\log_{10} p$ values versus the expectation under the null. Black points represent the complete meta-analysis, with a substantial departure from the null at the tail (values > 8 are represented along the top of the plot as triangles). Dark blue points show the distribution after removing 11 previously published loci, demonstrating a still notable excess. Light blue points show the distribution after removing all 40 loci which replicate at least nominally. In all the cases the overall distribution is marginally inflated ($\lambda_{GC} < 1.16$).

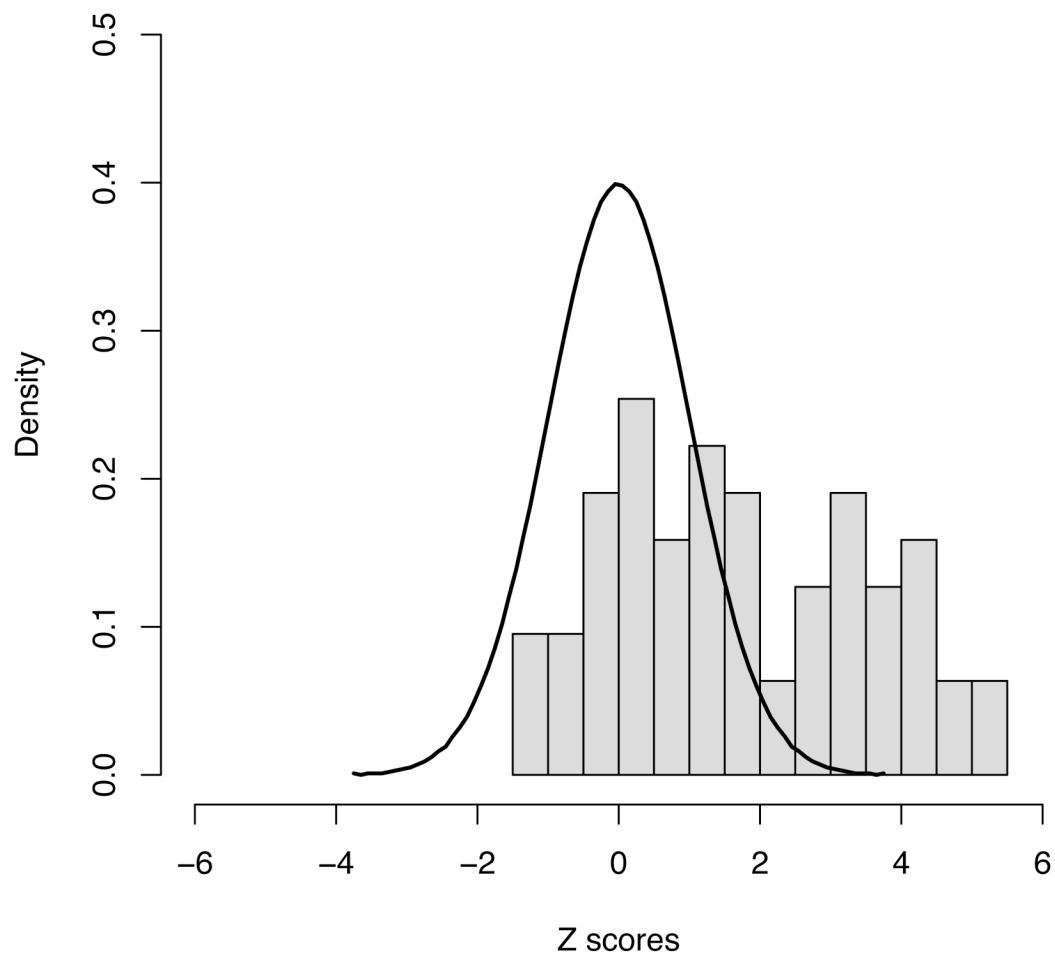


Figure 3.

Distribution of observed Z scores from the 63 novel regions explored, along with the expected distribution under the null (a standard normal with mean 0 and variance 1). Even setting aside the 21 regions reaching genome-wide significance, the distribution is highly skewed – 4 more results exceed a Z of 2 (1 would be expected by chance under the null) whilst none showed a Z of less than -2 (same expectation under the null) suggesting that even more of the regions investigated here are likely to constitute true positive associations when additional data become available.

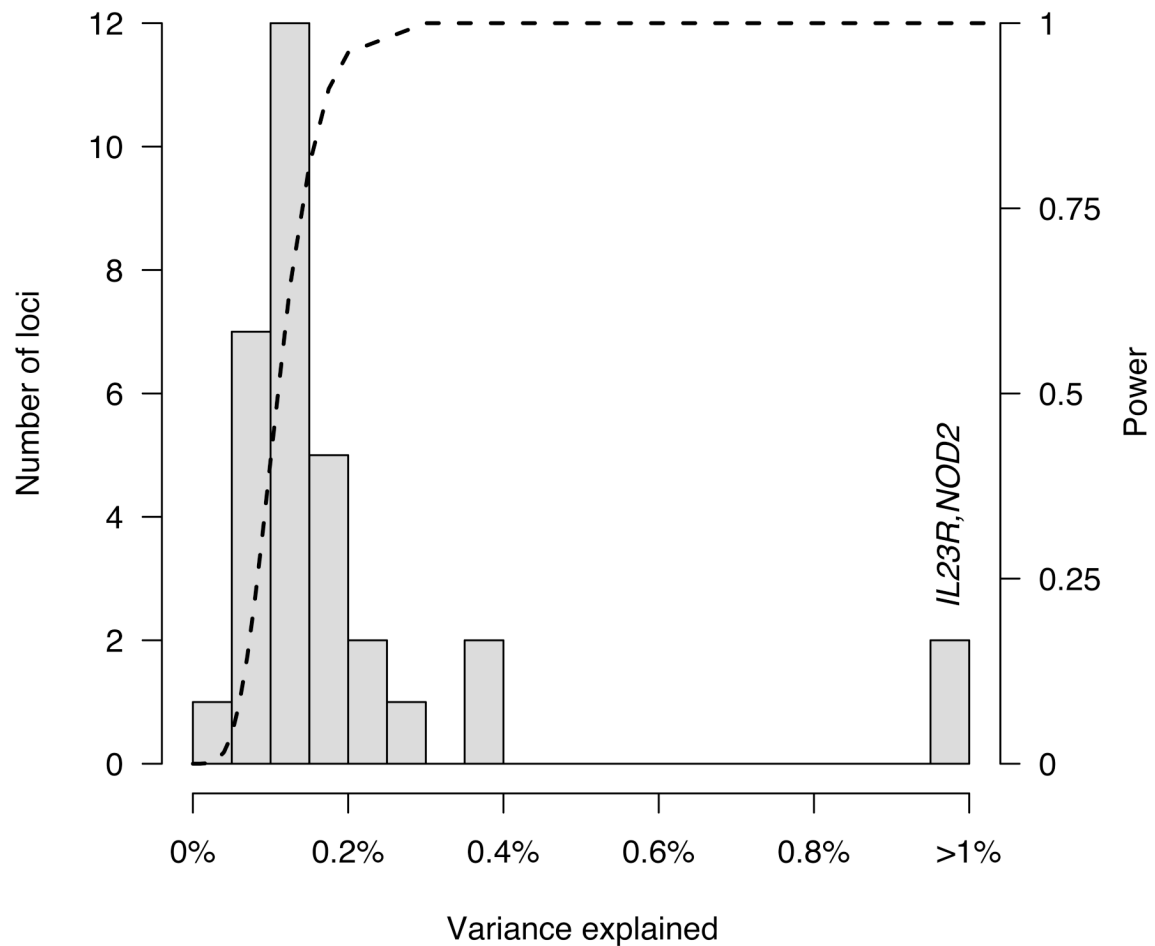


Figure 4.

Histogram of percent variance explained by each of the 32 established CD risk loci. The distribution resembles the long postulated exponential distribution of effect sizes. Dashed line shows the joint power for our meta-analysis to detect ($p < 5 \times 10^{-5}$), and for our replication sample to replicate (at Bonferroni corrected p values), a 20% variant explaining a given fraction of variance. Note how quickly this curve moves from nearly zero power to detect tiny effects (less than one tenth of one percent) to nearly full power to detect larger effects (presuming they are well covered by the current generation of GWAS chips). Complete power near the origin would likely reveal a more complete exponential distribution, with many very small effects. These are likely to increase somewhat once the causal variant or variants are identified in each locus. Indeed, *NOD2* and *IL23R* are distant outliers, each explaining 1-2% of total variance, partially because multiple causal variants have already been discovered at these loci^{6,13}.

Table 1**Samples used (post QC) in this study**

	NIDDK	BEL/FR	UKIBDGC	Total
Scan cases	946	536	1,748	3,230
Scan controls	977	914	2,938	4,829
Replication cases	0	1,082	1,243	2,325
Replication controls	0	787	1,022	1,809
Replication Trios	720	619	0	1,339
Nationality	USA/Canadian	Belgian/French	British	
Scan Platform	Illumina HumanHap300	Illumina HumanHap300	Affymetrix GeneChip 500K	
Replication Platform	Sequenom	Illumina GoldenGate	Sequenom	

Table 2
Convincingly (Bonferroni $p < 0.05$) replicated CD risk loci

SNP	Chr	Critical region	Scan	Replication	Combined	Num. genes	Gene of interest	RAF	Risk allele	Case Ctrl	Odds ratios	TDT
(a) Previously published loci												
rs111465804	1p31	67.4 *	1.01×10^{-35}	3.1×10^{-29}	6.66×10^{-63}	NA	<i>IL23R</i>	0.933	T	2.50	2.77	
rs3828309	2q37	230.9 *	1.13×10^{-20}	7.67×10^{-14}	2.36×10^{-32}	NA	<i>ATG16LI</i>	0.533	G	1.28	1.30	
rs3197999	3p21	48.73 - 49.87	2.16×10^{-7}	5.64×10^{-7}	1.15×10^{-12}	35	<i>MST150</i>	0.271	A	1.20	1.20	
rs4613763	5p13	40.32 - 40.48	4.52×10^{-22}	2.79×10^{-8}	6.82×10^{-27}	0	<i>PTGER4</i> **	0.125	C	1.32	1.28	
rs2188962	5q31	131.44 - 131.90	4.58×10^{-9}	3.52×10^{-11}	2.32×10^{-18}	7		0.425	T	1.25	1.26	
rs11747270	5q33	150.15 - 150.32	6.36×10^{-11}	2.57×10^{-7}	3.40×10^{-16}	3	<i>IRGM</i>	0.090	T	1.33	1.31	
rs4263839	9q32	114.61 - 114.78	3.92×10^{-7}	6.58×10^{-5}	2.60×10^{-10}	2	<i>TNFSF15</i>	0.677	G	1.22	1.07	
rs10995271	10q21	64.05 - 64.12	1.90×10^{-11}	1.61×10^{-10}	4.46×10^{-20}	1	<i>ZNF365</i>	0.387	C	1.25	1.53	
rs11190140	10q24	101.26 - 101.32	1.71×10^{-10}	1.69×10^{-7}	3.06×10^{-16}	1	<i>NKX2-3</i>	0.478	T	1.20	1.28	
rs2066847	16q12	49.3 *	NA	1.49×10^{-24}	2.98×10^{-24}	NA	<i>NOD2</i>	0.018	C	3.99	2.57	
rs2542151	18p11	12.73 - 12.88	1.19×10^{-11}	2.41×10^{-7}	5.10×10^{-17}	1	<i>P TPN2</i>	0.152	G	1.35	1.14	
(b) Novel loci												
rs2476601	1p13	113.79 - 114.17	1.81×10^{-5}	0.000101	1.46×10^{-8}	7	<i>P TPN22</i>	0.899	G	1.31	1.17	
rs2274910	1q23	157.65 - 157.72	3.50×10^{-7}	0.000481	1.46×10^{-9}	2	<i>ITLN1</i>	0.682	C	1.14	1.62	
rs9286879	1q24	169.54 - 169.67	4.02×10^{-7}	0.000321	1.53×10^{-9}	0		0.243	G	1.19	1.08	
rs11584383	1q32	197.60 - 197.77	6.82×10^{-7}	2.34×10^{-6}	1.43×10^{-11}	3		0.697	T	1.18	1.20	
rs10045431	5q33	158.69 - 158.76	8.80×10^{-9}	3.66×10^{-6}	3.86×10^{-13}	1	<i>IL12B</i>	0.708	C	1.11	1.36	
rs6908425	6p22	20.63 - 20.84	2.52×10^{-7}	0.000278	8.96×10^{-10}	1	<i>CDKALI</i>	0.780	C	1.21	1.09	
rs7746082	6q21	106.52 - 106.62	3.70×10^{-6}	7.7×10^{-6}	2.44×10^{-10}	0		0.289	C	1.17	1.19	
rs2301436	6q27	167.32 - 167.52	3.30×10^{-7}	3.26×10^{-7}	1.04×10^{-12}	3	<i>CCR6</i>	0.463	T	1.21	1.16	
rs1456893	7p12	50.03 - 50.11	4.92×10^{-5}	1.1×10^{-5}	4.60×10^{-9}	0		0.678	A	1.20	1.14	
rs151398	8q24	126.60 - 126.62	4.90×10^{-6}	0.000109	4.50×10^{-9}	0		0.619	A	1.08	1.25	
rs10758669	9p24	4.94 - 5.26	6.80×10^{-7}	0.00043	3.46×10^{-9}	3	<i>JAK2</i>	0.348	C	1.12	1.21	
rs17582416	10p11	35.30 - 35.60	8.48×10^{-6}	2.53×10^{-5}	1.79×10^{-9}	3		0.345	G	1.16	1.26	
rs7927894	11q13	75.80 - 76.02	1.43×10^{-7}	0.000732	1.32×10^{-9}	1	<i>C11orf30</i>	0.386	T	1.16	1.07	
rs11175593	12q12	38.61 - 39.31	1.33×10^{-7}	0.000165	3.08×10^{-10}	3	<i>LRRK2, MUC19</i>	0.017	T	1.54	1.44	
rs3764147	13q14	43.13 - 43.54	1.61×10^{-7}	1.33×10^{-7}	2.08×10^{-13}	3		0.221	G	1.25	1.19	
rs2872507	17q21	34.63 - 35.34	2.12×10^{-6}	0.000292	5.00×10^{-9}	17	<i>ORMDL3</i>	0.473	A	1.12	1.24	
rs744166	17q21	37.74 - 37.95	5.94×10^{-6}	9.15×10^{-8}	6.82×10^{-12}	4	<i>STAT3</i>	0.565	A	1.18	1.25	
rs1736135	21q21	15.73 - 15.76	2.06×10^{-5}	4.58×10^{-5}	7.40×10^{-9}	0		0.565	T	1.18	1.10	
rs762421	21q22	44.43 - 44.48	1.08×10^{-5}	1.59×10^{-5}	1.41×10^{-9}	1	<i>ICOSLG</i>	0.389	G	1.13	1.21	

RAF is risk allele frequency in control samples (see Supplementary Table 5 for details). Critical region is in NCBI B35 coordinates, with definition as described in Methods. Risk alleles are defined relative to the + strand of the reference.

* regions where causal variants have been convincingly mapped, rendering the LD window uninformative.

** *PTGER4* is outside the critical region, but was implicated via eQTL analysis.

Table 3

Nominally ($p < 0.05$) replicated CD risk loci

SNP	Chr	Critical region	Scan	p values		Num. genes	Gene of interest	RAF	Risk allele	Odds ratios	
				Replication	Combined					Case Ctrl	TDT
rs4807569	19p13	1.05 - 1.15	1.16×10^{-8}	0.00347	2.12×10^{-9}	2	<i>GCKR</i>	0.217	C	1.02	1.26
rs780094	2p23	27.30 - 27.77	3.82×10^{-6}	0.00381	3.14×10^{-7}	22		0.397	T	1.08	1.13
rs3763313	6p21	32.44-32.79 *	1.45×10^{-8}	0.00602	5.20×10^{-9}	7	<i>BTNL2, DRA, DRB, DQA</i>	0.188	C	1.19	1.01
rs13003464	2p16	61.09 - 61.14	3.44×10^{-5}	0.00565	4.60×10^{-6}	1		0.376	G	1.16	1.08
rs991804	17q12	29.57 - 29.70	4.02×10^{-6}	0.0135	1.07×10^{-6}	4	<i>CCL2, CCL7</i>	0.726	C	1.1	1.08
rs12529198	6p25	5.04 - 5.11	7.08×10^{-7}	0.0192	6.96×10^{-7}	1		0.062	G	1.12	1.19
rs17309827	6p25	3.36 - 3.42	2.08×10^{-6}	0.0391	2.74×10^{-6}	1	<i>LYRM4</i>	0.639	T	1.1	1.02
rs7758080	6q25	149.54 - 149.65	7.28×10^{-6}	0.044	8.78×10^{-6}	0		0.274	G	1.12	0.99
rs8098673	18q11	17.74 - 17.93	3.18×10^{-5}	0.0443	2.88×10^{-5}	0	<i>SLC22A23</i>	0.329	C	1.05	1.09
rs917997	2q11	102.31 - 102.64	2.16×10^{-5}	0.0493	2.22×10^{-5}	5		0.222	T	1.05	1.11

RAF is risk allele frequency in control samples (see Supplementary Table 5 for details). Critical region is in NCBI B35 coordinates, with definition as described in Methods. Risk alleles are defined relative to the + strand of the reference.

* SNPs with $p < 0.0001$ were observed throughout the MHC from 30.2 – 32.9 Mb but only this largest signal from the region was followed up. More detailed study of the MHC will be required to identify and localize potentially independent signals from this region.